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- (4) 1,3-Oxathiolenes useful in the treatment of hepetitis.
- The present invention relates to the use of nucleoside enalogues in the treatment of viral infections.

 More specifically it is concerned with the use of 1,3-axistricians nucleoside enalogues in the treatment of hepatitis, in particular hepatitis 8.

EP 0 515 144 A1

The present invention relates to the use of nucleoside enalogues in the treatment of viral infections. More specifically it is concerned with the use of 1,3-oxistholene nucleoside enalogues in the treatment of hepatits, in particular hepatitis 8.

Hepatitis 8 is a viral disease transmitted orally or parenterally by contaminated material such as blood and blood products, contaminated needles, sexually and vertically from infected or carrier mothers to their off-spring. In those areas of the world where the disease is common, vertical transmission at an early age results in a high proportion of infected individuals becoming chronic carriers of hepatitis 8. There are an estimated 280,000,000 carriers of hepatitis 8 worldwide. At the present time there are no effective chemotherspectic agents for the treatment of hepatitis 8 infections.

European patent publication 0382526A describes a series of 1,3-exathibians nucleoside analogues having antiviral activity, in particular activity against HIV, the causative agent of AIDS. We have now found that certain J of the compounds described in EP 0382526A are active both in vitro and in vivo against the hepatitis 8 virus.

The invention accordingly provides, in a first sepect, a method for the treatment of an animal, including man, infected with the hepatitle 5 virus comprising the administration of an effective amount of a compound of formula (I) or a pharmacoutically acceptable derivative thereof

wherein R₁ is hydrogen or an acyt;

R₂ is a purine or pyrimidine base or an analogue or derivative thereof;

Z is S. S=0 or SO;

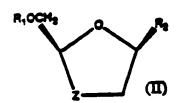
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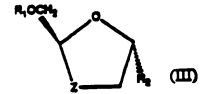
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provided that R_2 is not dyscalne-ligher, the compound of formula (I) is in the ris configuration. R_1 is hydrogen and Z is S. I

It will be appreciated by those skilled in the art that the compounds of formula (f) contain at least two chiral centres (shown as * in formula (f)) and thus exist in the form of two pairs of optical isomers (i.e. enantiomers) and mixtures thereof including recentle mixtures. Thus the compounds of formula (f) may be either ofe isomers, as represented by formula (ff), or mixtures thereof. Each of the cis and trans isomers can exist as nos of two enantiomers as a mixtures thereof including canonic mixtures. All such isomers and mixtures thereof including recentle mixtures are included within the scope of the invention:





The compounds of formula (I) are preferably in the form of their cle isomers.

It will also be appreciated that when Z is \$=0 the compounds exist in two additional recemic forms as shown in formulas (IIs) and (IIb) which differ in the configuration of the oxide oxygen atom relative to the 2,5-substituents. The compounds of the invention additionally embrace such isomers and mixtures thereof.

. The purine or pyrimidine base $R_{\rm 2}$ will be linked at the 9- or 1- position respectively.

By purine or pyrimidine base or an analogue thereof is meant a purine or pyrimidine base found in nucleosides or an analogue thereof which mimics such bases in that their structures (the kinds of atoms and their arrangement) are similar to the normal bases but may either possess additional or lack certain of the functional properties of the normal bases. Such analogues include those derived by replacement of a CH₂ molety by a nitrogen atom (for example, 5-exapyrimidines such as 5-exacytosine) or vice verse (for example 7-deazapurines, for example 7-deazadenosine or 7-deazaguanosine) or both (e.g. 7-deazadenosine or 7-deazaguanosine) or both (e.g. 7-deazadenosine or 7-deazaguanosine) or both (e.g. 7-deaza, 8-exapurines). By derivatives of such bases or analogues are meant those compounds wherein ring substituents are either incorporated, removed or modified by conventional substituents known in the art e.g. halogen, hydroxyl, amino, C₁₋₆ alkyl. Such purine or pyrimidine bases, analogues and derivatives will be well known to those skilled in the art.

Conveniently the group R2 is selected from:

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wherein Rije selected from the group consisting of: hydrigin and C₁₋₀ alicyl, unsubstituted or substituted with a heterostom;

Rayand Re are independently selected from the group consisting of: hyperioges. Che sinyl, bramine, chlorine, seede, and lodine;

R_i is selected from the group consisting of: hydrogen, CN, carboxy, ethoxycarboxyl, carbamoyl and thiocarbamoyl; and

X and Y are independently selected from the group consisting of: bromine, chlorine, fluorine, iodine, amino and hydroxy groups.

Preferebly Re 104

wherein Rs and Rs are se defined hereinabove.

Z is preferably -S-.

R₁ and R₄ are preferably hydrogen or C₁₋₂ alkyl.

Ra is preferably CH2 or F.

X and Y are preferably both NH2

It will be appreciated by one of skill in the art that when R₁ is an acyl group, the compounds of formula (I) are esters. Preferred esters include a carboxyl function R-CO-O in which the non-carbonyl molety R is selected from hydrogen, straight or branched chain alkyl (e.g. methyl, ethyl, n-propyl, t-butyl, n-butyl), alkoxyalkyl (e.g., methoxymethyl), arallyl (e.g. benzyl), aryloxyalkyl (e.g. phenoxymethyl), aryl (e.g. phenyl optionally substituted by hatogen, C1-4 sikyl or C1-4 sikoxy); substituted dihydro pyridinyl (e.g. N-methyldihydro pyridinyl); sulphonate esters such as alkyl- or aralkylsulphonyl (e.g. methanesulphonyl); sulfate esters, amino acid esters (e.g. L-valyl or L-isoleucyl) and mono-, di- or tri-phosphata estera.

Also included within the scope of such esters are esters derived from polyfunctional acids such as carbonylic acids containing more than one carboxyl group, for example, dicarboxylic acids HO₂C(CH₁)₂CO₂H where n is an integer of 1 to 10 (for example, succinic acid) or phosphoric acids. Methods for preparing such esters from the corresponding alcohol are well known. See, for example, Hahn et al., "Nucleotide Dimers as Anti Human Immunodeficiency Virus Agents', <u>Nucleotide Analogues</u>, pp. 158-158 (1989) and Busso et al., "Nucleotide Dimers Suppress HIV Expression in Vitro". AIDS Research and Human Retroviruses, 4(6), pp. 449-455 (1986).

With regard to the above described esters, unless otherwise specified, any alkyl molety present advantageously contains 1 to 16 cerbon atoms, perticularly 1 to 4 cerbon stoms and could contain one or more double bonds. Any anyt moiety present in such esters advantageously comprises a phenyl group.

In particular the esters may be a C_{t-10} sligt ester, an unsubstituted berzoyl ester or a berzoyl ester substituted by at least one halogen (bromine, chlorine, fluorine or lodine), C₁₋₄ alityl, saturated or unesturated C1-4 alkaxy, nitro or trifluoromethyl groups.

By the term "pharmaceutically acceptable derivative" is meent any pharmaceutically acceptable sait of a compound of formula (I) or any other compound which, upon administration to the recipient, is capable of providing (directly or indirectly) a compound of formula (I) or an antivirally active metabolite or residue thereof.

It will be appreciated by those sidled in the art that the compounds of formula (f) may be modified to provide pharmaceutically acceptable derivatives thereof, at functional groups in both the base moisty and at the R. group of the exactilolane ring. Modification at all such functional groups are included within the ecope of the invention.

Pharmaceutically acceptable saits of the compounds of formula (I) include those derived from pharmaceutiically ecceptable inorganic and organic acids and bases. Examples of suitable acids include hydrochloric, hydrobromia, sulphurle, nitrie, perchlorie, fumeric, meleic, phosphorie, glycollie, lactic, sellcytie, succinie, toluenep-sulphonic, tartaric, acetic, citric, methanesulphonic, formic, berzolc, malonic, naphthelene-2-sulphonic and benzenesulphonic acids. Other acids such as oxalic, while not in themselves pharmaceutically acceptable, may be useful in the preparation of saits useful as intermediates in obtaining the compounds of the invention and their pharmaceutically acceptable acid addition saits.

Salts derived from appropriate bases include alkali metal (e.g., sodium), alkaline earth metal (e.g., magnesium), ammonium and NR4+ (where R is C1.4 alkyl) saits.

References hereinafter to a compound according to the invention includes both the compound of formula

(i) and its pharmacoutically acceptable derivatives. Specific compounds of formula (I) include:

trans-2-hydroxymethyl-5-(cytosin-1'-yl)-1,3-exathiolane;

cis-2-benzoyloxymethyl-5-(cytosin-1'-yl)-1,3-oxethiclene, trans-2-benzoyloxymethyl-5-(cytosin-1'-yl)-

1,3-examinisme, and mixtures thereof;

c/s=2-hydroxymethyt-5-(N₄'-ecetyt-cytosin-1'-yt)-1,3-oxathiclene, /rata-2-hydroxymethyt-5-(N₄'-ecetyt-

cytosin-1'-yl)-1.3-exathiclane, and mixtures thereof;

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cis-2-benzoyloxymethyl-5-(N₄'-acetyl-cytosin-1'-yl)-1,3-oxathiolane, trans-2-benzoyloxymethyl-5-(N₄'acetyl-cytosin-1'-yl)-1,3-oxathiolane, and mixtures thereof;

cis-2 benzoyloxymethyl-5-(N₄'-ecetyl-5-fluorocytosin-1'-yl)-1,3-oxathiolane. trans-2 benzoyl-oxymethyi-5-(N_4 '-acetyi-5-fluorocytosin-1'-yi)-1,3-oxsthiolane, and mixtures thereof;

c/s-2-hydroxymethyl-5-(5'-fluorocytosin-1'-yl)-1,3-oxethiclene. trans-2-hydroxymethyl-5-(5'-fluorocytosin-1'-yi)-1,3-oxatholane, and motures thereof.

- cis-2-hydroxymethyl-5-(cytosin-1'-yl)-3-oxo-1,3-oxathiolane:
 - cis-2-hydroxymethyl-5-(thymin-N-1'-yl)-1,3-oxathidane; and
 - cis-2-hydroxymethyl-5-(N,N-dimethyleminomethylenecytosin-1'-yl)-1,3-oxethiolene;
- in the form of a recemic mixture or a single ensythmer.

The compounds of formula (I) are preferably in the form of the cis compounds and contain two chiral centres (shown in formula (i) by *).

The compound of formula (I) is preferably in the form of a recemic modure or a single enantiomer but a mixture of enantiomers in any ratio may be employed in the invention. Most preferably, the compound of formula: (I) is in the form of its (-) enantiomer.

The compounds of formula (I) and their individual enandomers may be prepared by any method known in the art for the preparation of compounds of analogous structure for example by the methods described in European patent publication 0382526A.

In a further or atternative aspect there is provided a compound of formula (I) as defined hereinsbove or a pharmaceutically acceptable derivative thereof for use in the manufacture of a medicament for the treatment of hepatitis 8.

As will be appreciated by those skilled in the art, references herein to treatment extend to prophylaxis as well as to the treatment of established infections of symptoms.

The compounds of formula (I) both as the recemic moture and as the individual enentioners have been found to inhibit the hepatite 5 virus both in vitro and in vivo.

It will be appreciated that the amount of a compound of the invention required for use in treatment will vary not only with the particular compound selected but also with the route of edministration, the nature of the condition being treated and the age and condition of the patient and will be ultimately at the discretion of the attendant physician or veterinarian. In general however a suitable dose will be in the range of from about 0.1 to about 750 mg/kg of bodyweight per day preferably in the range of 0.5 to 60 mg/kg/day, most preferably in the range of 1 to 20 mg/kg/day.

The desired dose may conveniently be presented in a single dose or as divided doses administered at appropriete intervals, for example as two, three, four or more sub-doses per day.

The compound is conveniently administered in unit dosage form; for example containing 10 to 1500 mg. conveniently 20 to 1000 mg, most conveniently 50 to 700 mg of eative ingredient per unit dosage form.

Ideally the active ingredient should be administered to achieve peak pleams concentrations of the active compound of from about 1 to about 75 µM, preferably about 2 to 50 µM, most preferably about 3 to about 30 μM. This may be achieved, for example, by the intrevenous injection of a 0.1 to 5% solution of the active ingredient, optionally in sellne, or orally administered as a bolus containing about 1 to about 100 mg of the active ingredient. Desirable blood levels may be maintained by a continuous infusion to provide about 0.01 to about 5.0 mg/tg/hour or by intermittent infusions containing about 0.4 to about 15 mg/tg of the ective ingredient.

While it is possible that, for use in therapy, a compound of the invention may be administered as the raw chemical it is preferable to present the active ingredient as a phermacoutical formulation.

The invention thus further provides a pharmacoutical formulation comprising a compound of formula (I) or a pharmaceutically acceptable derivative thereof together with one or more pharmaceutically acceptable carriers therefor and, optionally, other therepeutic and/or prophylactic ingredients. The certier(s) must be "acceptable" in the sense of being compatible with the other ingredients of the formulation and not deletarious to the recipient thereof.

Pharmaceutical formulations include those suitable for oral, rectal, nasal, topical (including buccal and sublingual), veginal or parentaral (including intramuscular, sub-outaneous and intravenous) administration or in a form suitable for administration by inhalation or insuffiction. The formulations may, where appropriate, be conveniently presented in discrete dosege units and may be prepared by any of the methods well known in the art of pharmacy. All methods include the step of bringing into association the active compound with liquid carriers or finely divided solid carriers or both and then, if necessary, shaping the product into the desired formulation.

Pharmaceutical formulations suitable for oral edministration may conveniently be presented as discrete units such as capsules, cachets or tablets each containing a predetermined amount of the active ingredient; as a powder or granules: as a solution, a suspension or as an emutaion. The active ingredient may also be

presented as a bolus, electuary or pasts. Tablets and capsules for oral administration may contain conventional excipients such as binding agents, fillers, lubricants, disintegrants, or wetting agents. The tablets may be coated according to methods well known in the art. Oral liquid preparations may be in the form of, for example, aqueous or only suspensions, solutions, emulsions, syrups or elixirs, or may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may contain conventional additives such as suspending agents, emulsifying agents, non-equeous vehicles (which may include edible oils), or preservatives.

The compounds according to the invention may also be formulated for parenteral administration (e.g. by injection, for example bolus injection or continuous infusion) and may be presented in unit dose form in ampoules, pre-filled syringes, small volume infusion or in multi-dose containers with an added preservative. The compositions may take such forms as suspensions, solutions, or emulsions in oily or equeous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredient may be in powder form, obtained by aseptic isolation of startle solid or by lyophilization from solution, for constitution with a suitable vehicle, e.g. startle, pyrogen-free water, before use.

For topical administration to the epidermis the compounds according to the invention may be formulated as cintments, creams or lodons, or as a transformal patch. Cintments and creams may, for example, be formulated with an equeous or only base with the addition of suitable thickening and/or gelling agents. Lotons may be formulated with an equeous or only base and will in general also contain one or more emulaifying agents, stabilizing agents, dispersing agents, suspending agents, thickening agents, or coloring agents.

Formulations suitable for topical administration in the mouth include lozanges comprising active ingredient in a flavored base, usually sucrose and acadis or gum tragacenth; pastiles comprising the active ingredient in an inert base such as gelatin and glycerin or sucrose and acadis; and mouthwashes comprising the active ingredient in a suitable liquid carrier.

Pharmaceutical formulations suitable for rectal administration wherein the carrier is a solid are most preferably presented as unit dose suppositories. Suitable carriers include cocce butter and other materials commonly used in the erc. and the suppositories may be conveniently formed by administure of the active compound with the softened or method carrier(s) followed by chilling and shaping in moulds.

Formulations suitable for veginal administration may be presented as pessaries, tempons, creams, gais, pastes, foams or sprays containing in addition to the active ingredient such centers as are known in the art to be appropriate.

For intra-nasal administration the compounds of the invention may be used as a liquid apray or dispersible powder or in the form of drops.

Orope may be formulated with an equeous or non-equeous base also comprising one or more dispersing agents, solubilizing agents or suspending agents. Uquid spraye are conveniently delivered from pressurized nacks.

For administration by inhelation the compounds according to the invention are conveniently delivered from an insuffictor, nebulizar or a pressurized pack or other convenient means of delivering an aerosal spray. Pressurized packs may comprise a suitable propellant such as dichlorodifluoromethene, trichloroduoromethene, dictionostrafluorosthane, carbon dioxide, nitrogen or other suitable gas. In the case of a pressurized serosal the desage unit may be determined by providing a valve to deliver a metered amount.

Alternatively, for administration by inhalation or insuffiction, the compounds according to the invention may take the form of a dry powder composition, for example a powder mix of the compound and a suitable powder base such as factose or starch. The powder composition may be presented in unit desage form in, for example, capsules or cartridges or e.g. gelatin or bi-size packs from which the powder may be administrated with the aid of an inhalator or insuffictor.

When desired the above described formulations adapted to give sustained release of the active ingredient may be employed.

The pharmaceutical compositions according to the invention may also contain other active ingredients such as antimicrobial agents, or preservetives.

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The compounds of the invention may also be used in combination with other therapeutic agents for example other antiinfective agents. In particular the compounds of the invention may be employed together with known antiviral, antibacterial, antifungal or immunomodulating agents.

The invention thus provides, in a further aspect, a combination comprising a compound of formula (I) or a physiologically acceptable derivative thereof together with another therapeutically active agent, in particular an antiverse, antifungal or immunomodulating agents.

The combinations referred to above may conveniently be presented for use in the form of a pharmaceutical formulation and thus pharmaceutical formulations comprising a combination as defined above together with a pharmaceutically acceptable carrier therefor comprise a further aspect of the invention.

The individual components of such combinations may be administered either sequentially or simultaneously in separate or combined pharmacoutical formulations.

When a compound of formula (i) or a pharmaceutically acceptable derivative thereof is used in combination with a second therapeutic agent active against the same virus the dose of each composed may be either the same as or differ from that when the compound is used alone. Appropriate doses will be readily appreciated by those skilled in the art.

The invention is illustrated by the following examples which should not be interpreted as a limitation of the invention.

Example 1

Cis- and trans-2-benzoyloxymethyl-5-(Na'-acetyl-5'-fluoro-cytosin-1'-yl)-1,3-oxathiolane

5-Fluorocytosine (4.30 g, 33.3 mmol), hexamethyldisilezane (25 ml) and arranonium suifate (120 mg) were boiled under reflux until the cytosine dissolved (3 hours) and then further refluxed for 2 hours. The hexamethyldisilezane is evaporated in vacuo and toluene (100 ml) was added to the residue to co-evaporate the solvents. The resulting solution, bis(trimethylaskyl)-fluoro-cytosine in dichloromethane (40 ml) was added under arronned as solution of 2-benzoyloxymethyl-5-acetony-1,3-oxachiolane (8.537 g, 30.3 mmol) in dry dichloromethane (100 ml) and molecular sieves (4A, 2 g) previously prepared under argon and cooled at O°C for 20 minutes. ([Trifluoromethane-cultonyl)oxyltrimethylsilane (6 ml, 31 mmol) was added to this mixture at O°C and the utes. ([Trifluoromethane-cultonyl)oxyltrimethylsilane (6 ml, 31 mmol) was added to this mixture was then treated with resulting solution was stirred at 25°C for approximately 18 hours. The reaction mixture was then treated with resulting solution was stirred at 25°C for approximately 18 hours. The reaction mixture was then treated with resulting solution was stirred at 25°C for approximately 18 hours. The reaction mixture was then treated with resulting solution of sodium bicarbonate and stirred at room temperature for 2 hours. The filtrate was shaken two times with 300 ml of brine and one time with distilled water. The organic layer was dried over magnetium sulfate, filtered and evaporated to dryness. This afforded a crude 5-fluoro-cytosine derivative (10.1 magnetium sulfate, filtered and evaporated to dryness. This afforded a crude 5-fluoro-cytosine derivative (10.1 magnetium sulfate, filtered and evaporated to dryness.

g). R.O.57 (EtOAc:MeCH 9:1).

This residue was acetylated in the next step without further purification. The crude material was dissolved in the residue was acetylated in the next step without further purification. The flask was then immersed in and dimethyl artinopyridine (120 ml) in a 500 ml round bottom flask under argon. Triethylamine (12.7 ml, 91.9 mmol) and dimethyl artinopyridine (111 mg, 0.9 mmol) were added to the solution. The flask was then immersed in an ice bath for 1 hour under argon. Acede anhydride (4.3 ml, 45 mmol), distilled over sodium acestate, was syninged into the cooled flask. The mixture was stirred overright and then carefully decembed with distilled water followed into the cooled flask. The mixture was solution. The product was then weathed with distilled water followed by brine solution. The methylene chloride portions were dried and evaporated under high vacuum to drylowed by brine solution. The methylene chloride portions were dried and evaporated under high vacuum to drylowed by brine solution. The methylene as a coloriese form, weighing 9.8 g after drying. Flash chromatography of this metarial using edhylacestatemethanol (9:1) afforded 3.1 g. 7.8 mmol (46%) pure trans-(berzoyloxymethyl-5-(N₄'-acetyl-5'-fluoro-cytosin-1'-yl)-1,3-oxistholane).

trans-isomer: R.O.65 in ethyl acetate:methanol 9:1

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U.V.: (MeOH) Lambda mez: 309 nm
  'H-NMR & (ppm in CDCI)
         8.77 (b. 1H; C.'-NH-AC)
          8.06 (m, 21t aromatic)
          7.70 (d. 11t Co'-15 Japon 344)
          7.62 (m. 1H; aromatic)
          7.49 (m. 21t; aromatic)
          8.51 (dd. 1H; C+11)
          5.91 (dd. 1H; CrH)
           4.48 (64. 2H; C.C.HOCOC.H.)
           3.66 (dd. 1H; C.-H)
           3.34 (dd, 1H; C4-H)
           2.58 (s. 3H; NH-COC<u>Hs</u>)
           cis-somer: R.O.58 in ethyl acetate:methanol 9:1
     U.V.: (MeOH) Lambda max: 309nm
     'H-NMR & (ppm in CDCL)
           8.72 (b. 1H; C.'-NH-AC)
            8.06 (m. 2)+; aromatic)
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            7.57 (d. 1H; Ca'-H, Ja-6.2Hz)
            7.60 (m. 1H; aromatic).
            7.49 (m, 2H; aromastc)
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. 6.32 (dd. 1H; C<sub>F</sub>H)
5.47 (dd. 1H; C<sub>F</sub>H)
4.73 (dd. 2H; C<sub>F</sub>CH<sub>2</sub>OCOC<sub>6</sub>H<sub>6</sub>)
3.62 (dd. 1H; C<sub>6</sub>H)
3.19 (dd. 1H; C<sub>6</sub>H)
2.55 (s. 3H; NH-COCH<sub>2</sub>)
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Example 2

Cla- and trans-hydroxymethyt-5-(5'-fluorocytosin-1'-yl)-1,3-oxethiclane

1.0 g (2.54 mmol) of trans-2-benzoyloxymethyl -5-(N₄'-ecstyl-5'-fluorocytosin-1'-yl)-1.3-oxethiolane was stirred in 25 ml of methanolic ammonia at 0° for 1 hour and then overnight at room temperature. The mixture was evaporated under reduced pressure. The residue as titurated twice (2 x 30 ml) with anhydrous ether. The solid residue was recrystalized in absolute ethanol to give 484 mg (1.95 mmol, 77%) of desired product trans-(hydroxymethyl-5-(5'-fluorocytosin-1'-yl)-1,3-oxethiolane); m.p. 219-221°C; R,=0.21 in ethyl acetate; methanol (9:1), which was identified by ¹H, ¹²C-NMR and U.V. Lambda max (H₂O) 280.9 nm.

1.2 g (3.05 mmol) of cis-2-benzoyloxymethyl-5-(N₄'-scetyl-6'-fluoro-cytosin-1'-yl)-1,3-axathiolane was stirred in 30 ml of methanolic ammonia at 0°C for 1 hour and then overnight at room temperature. The mixture was evaporated under reduced pressure. The residue was triturated twice (2 x 30 ml) with anhydrous ether. The solid residue was recrystalitzed in absolute ethanol to give 655 mg (2.64 mmol, 87%) of pure product cla-(hydroxymethyl-5-(5'-fluorocytosin-1'-yl)-1,3-axathiolane): m.p. 204-208°C; R,=0.21 in ethylacetata: methanol (9:1). The desired compound was identified by ¹H, ¹³C-NMR and U.V. Lambda max (H₂O) 280.9 nm. trans-isomer:

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C_2' C_4' C_5' C_6' C_6' 153.47 158.20 134.65 126.24 C_7 = 13.2 \text{ Hz} C_7 = 26.2 \text{ Hz}
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C₅ C₄ C₂ CH₂OH 88.20 36.18 87.16 64.71

de-isamen:

'H-NMR & (ppm in DMSO-du):

8.22 (d. 1H; C₄'-H, J₆-7.28 Hz)

7.843 (d. 2H; C₄'-NH₃)

8.16 (t. 1H; C₇-H)

5.43 (t. 1H; C₇-H)

5.19 (t. 1H; C₇-H)

3.77 (m. 2H; C₇-H₂OH)

3.35 (dd. 1H; C₄-H)

3.12 (dd. 1H; C₄-H)

.

c, '	c , '	c _s '	C61
153.46	158.14	134.63	126.32
	(² J _{CF} =14.0 Hz)	(J _{CF} =24.1 Hz)	(J _{CF} =32.5 Hz)
c,	c ₄	c ₂	CH ³ OH
86.82	36.80	86.77	62.32

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Example 3

Blological Results

(A) Newborn ducklings were infected with duck hepatitie 5 virus (DHSV). After 5 to 7 days post-infection, samples of blood were taken from the ducidings and examined for DHSV DNA using dot hybridization with a specific DNA probe (Mason et al., Proc. Netl. Acad. Sci. USA 79, pp. 3997-4001 (1982)). The livers were removed from dot-blot positive ducklings and used to produce primary hapatocyte cultures infected with DHBV as previously described (Tuttleman et al., J. of Virology, 58, pg. 17-25). After 2 days in culture, antiviral egents were added to the culture media. The media were changed every 2 days and at selected times, the calls were removed and the total DNA extracted.

The DNA was spotted on nitrocellulose paper and probed with the PP-labelled DHSV DNA probe in access. dance with the following procedure. The DNA from DHEV-infected hepetocytes was extracted and spotted onto a nitrocellulose filter. The above described ***P-nick translated-DHEV DNA (pDH-010 = DHEV) probe was used. The DNA was extracted from 6-cm cell culture dishes at various times post-plating, in the virus control (VC) group, cells were harvested at 2, 8, 8, 10, 14, 18 and 20 days. Duplicate samples were spotted for days 14, 18 and 20. In drug-treated groups, cells were harvested on days 8, 14 and 20. Drugs were added to the culture at 2 days post-pletting and maintained throughout media changes every 2 days. The total intracellular DNA was extracted from cells using the standard phenol extraction method. The cells in a 6-cm diameter Petri dish (approximately 5 x 10° cells) were lysed in a lysis buffer containing 0.2% SDS, 150 mM Tris-HCl pH 8.0, 10 mM EDTA, 5 mM EGTA, and 150 mM NeCt. The cell lysets was digested with 0.5 mg/ms of process E (available from Sigme) at 37°C for 2 hours and proteinized by extraction with an equal volume of phenol saturated with 20 mM Trie-HCl, pH 7.5, 0.5 mM EDTA and 0.1% 8-hydroxyquinoline. Concentrated ammonium ecetate (pH 7.0 (2.5 M)) was added to the equeous phase to yield a 0.25 M ammonium acetate solution and the nucleis acids were precipitated with 2 volumes of 100% ethanol. The pellet of nucleic acid was washed with ethanol and dried. The DNA was dissolved in a solution containing 12.5 mM Tris-HCL pH 7.5, 10 mM EDTA, 30% glycerol and 0.01% bramophenol blue. One breith of the DNA sample was spotted onto the narocelulose for dolblot analysis.

The drugs tested were scored on a scale of 0 (no activity) to ++++ (high activity).

The compounds tested were 1,3 existhiplenes and two known inhibitors of hepetitis 8, 2, 3-dideoxy-gusnomine (ddG) and 2.6-diaminopurine-9-8-0-2'.3'-dideoxyribofuranceide (ddDAPR)-(European Petent publicaton 0302780AL

The results are shown in Table 1.

Table 1

5	Compound	Activity
	trans-2-hydroxymethyl-5-(5'-fluorocytosin-1'	
10	-yl)-1,3-oxathiolane	+
	cis-2-hydroxymethyl-5-(5'-fluorocytosin-1'	
15	-yl)-1,3-oxathiolane	+++
13	cis-2-hydroxymethyl-5-(thymin-K-1'-yl)-	
	1,3-oxathiolane	++
20	cis-2-hydroxymethyl-5-(N,N-dimethylamino-	
	methylene cytosin-1'-yl)-1,3-oxathiolane	++++
25	ddG	++++
39	ddDAPR	****

Claims

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1. Use of a compound of formula (I) or a pharmacoutically acceptable derivative thereof in the manufacture of a medicament for the treatment of a hepatitis 8 infection:

wherein R₁ is hydrogen or an acyl;

 $\ensuremath{R_{2}}$ is a purine or pyrimidine base or an analogue or derivative thereof, and

Z iS S, S=0, or SO;

provided that R_2 is not cytosine when the compound of formula (1) is in the cir configuration, R_1 is hydrogen and Z is S.

2. The use according to claim 1, wherein the ester is selected from the group consisting of: R-CO-O-, wherein R is selected from hydrogen, straight or branched sikyl, sikoxysikyl, sraikyl, sryloxysikyl, sryl, and substituted dihydropyridinyl; sulphonate esters; sulfate esters; amino acid esters; mono- di- or tri-phosphate esters; esters of polyfunctional acids; and esters of phosphoric acids.

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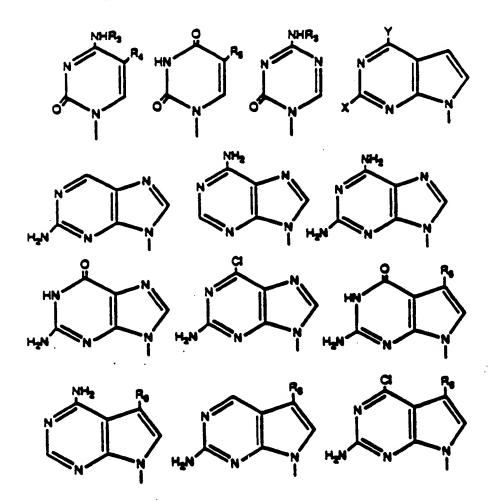
3. The use according to claim 1, wherein Z is S.

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4. The use eccording to claim 1, wherein R_2 of formula (I) is selected from the group consisting of:



wherein R_0 is selected from the group consisting of: hydrogen and C_{1-0} ellipt, unsubstituted or substituted with a heterostom;

R₄ and R₆ are independently selected from the group consisting of hydrogen, C₁₋₂ aliqu, bromine, chlorine, fluorine, and lodine;

Re is estected from the group consisting of: hydrogen, CN, carbony, ethonycarbonyl, carbamoyl and thiocerbamoyl; and

X and Y are independently selected from the group consisting of: bromine, chlorine, fluorine, lodine, amino and hydroxy groups.

S. The use eccording to claim 4, wherein R_{g} is

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wherein $R_{\rm 3}$ is selected from the group consisting of: hydrogen and $C_{\rm 1-0}$ alkyl unsubstituted or substituted with a heterostom: and

 R_4 is selected from the group consisting of: hydrogen, $C_{1-\delta}$ alkyl and bromine, chlorine, fluorine, and iodine.

- 6. The use according to claim 4 wherein R1 and R4 are hydrogen or C14 alkyl.
- 7. The use according to claim 4, wherein Re is CH2 or F.

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- ... S. The use according to claim 4, wherein X and Y are both NH₂.
 - The use according to claim 1, wherein the compound is selected from the group consisting of: trans-2-hydroxymethyl-5-(cytosin-1'-yl)-1,3-oxisthiolene;

cis-2-berzzykoxymethyl-5-(cytosin-1'-yl)-1,3-cxxthiclens, trans-2-berzzykoxymethyl-5-(cytosin-1'-yl)-1,3-cxxthiclens, and mbdures thereof;

1,3-examidiane, and modures thereof:

cis-2-hydroxymethyl-5-(N4'-ecstyl-cytosin-1'-yl)-1,3-examidians, trans-2-hydroxymethyl-5-(N4'-ecstyl-cytosin-1'-yl)-1,3-examidians, and modures thereof;

cis-2-benzoyloxymethyl-5-(N4'-acetylcytoein-1'-yl)-1,3-oxathiciene, trans-2-benzoyloxymethyl-5-(N4'-acetylcytoein-1'-yl)-1,3-oxathiciene and mixtures thereof;

cis-2-berzoylarymeth/-6-(N4'-ecetyl-6-fluorocytasin-1'-yi)-1,3-assittalene, trans-2-berzoy- larymethyl-... (N4'-ecetyl-6-fluoro-cytasin-1'-yi)-1,3-assittalene, end mbatares thereof;

ca-2-hydrograethyl-5-(5'-fluorocycein-1'-yl)-1,3-axethiolene, trene-2-hydrograethyl-5-(5'-fluorocycein-1'-yl)-1,3-axethiolene, trene-2-hydrograethyl-1-(5'-fluorocycein-1'-yl)-1,3-axethiolene, trene-2-hydrograethyl-1'-yl)-1,3-axethiolene, trene-2-hydrograethyl-1'-yl)-1,3-axethiolene, trene-2

cis-2-hydronymethyl-6-(gytosin-1'-yl)-3-axo-1,3-axethiclene;

cis-2-hydroxymethyl-5-(thymin-N-1'-yf)-1,3-costhiolene; and

cis-2-hydroxymethyl-5-(N,N-dimethyleminomethylene cytosin-1'-yl)-1,3-axisticlene; or pharmaceutically acceptable derivatives thereof, in the form of a recemic mixture or a single enemtiomer.

- 10. The use according to any one of claims 1 to 9, wherein the compound of formula (I) is present as a single enandomer or as a recemic mixture.
- 11. The use according to claim 10, wherein the compound of formule (I) is present as its (-) enantiomer.
- 12. The use eccording to claim 10, wherein the compound of formule (I) is present as its (+) enandomer.
- 13. The use according to any one of claims 1 to 9, wherein the compound is present in either its ole or trans configuration or mixture thereof.
 - 14. The use according to claim 13, wherein the compound of formula (I) is present in its de configuration.
- 15. Use of cie-2-hydroxymethyl-5-(5'-fluorocytosin-1'-yl)-1,3-oxethiclens in the menufacture of a medicament for the treatment of a hepatitis B infection.
 - 16. Use of cis-hydroxymethyl-5-(N,N-dimethylaminomethylene cytosin-1'yi)-1,3-exathicisne in the manufacture of a medicament for the treatment of hepatitis 8 infection.
- 17. The use according to any one of claims 1 to 9, wherein the medicament is edepted for oral, perenteral, rectal, need, veginal, or topical edministration.
 - 18. The use according to claim 17, wherein said medicament is edministered at a dose of about 0.1 to 750

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implies of bodyweight per day.

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- 19. The use according to claim 17, wherein said medicament is present in dosage unit form in the medicament.
- 20. The use according to claim 19, wherein the dosage unit form contains approximately 10 to 1500 mg of the compound of formula (I).
 - 21. The use according to any one of claims 1 to 9, wherein said medicament is administered with a pharmeceutically acceptable carrier.
- 22. The use according to any one of claims 1 to 7, wherein the medicament is edministrated in combination with a therapeutically active agent selected from the group consisting of: antiviral, antibacterial, antifungal and immunomodulating agents.

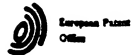


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